



UNIVERSITI PUTRA MALAYSIA

**IMPROVEMENT OF MOLECULAR METHODS FOR
DETECTION OF PATHOGENIC ESCHERICHIA COLI**

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IMPROVEMENT OF MOLECULAR METHODS FOR DETECTION OF PATHOGENIC *ESCHERICHIA COLI*

By

NAGI AHMED ABDULLAH AL-HAJ

**Thesis Submitted to the School of Graduate Studies, Universiti Putra Malaysia in
Fulfilment of the Requirement for the Degree of Doctor Philosophy**

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DEDICATION

To my mother who patiently awaited my return to Yemen after the completion of my studies but unfortunately she passed away. May Allah blessed her and accepted her in the high paradise with my father.

To my wife and our children, Ahmed, Sala and Hamzah. They provided a much needed balance in my life and were much more reasonable about the many, many lost weekends than I could possibly have hoped for. This dissertation is dedicated to them .

Abstract of thesis presented to the Senate of Universiti Putra Malaysia in fulfillment of the requirement for the degree of Doctor of Philosophy

**IMPROVEMENT OF MOLECULAR METHODS FOR DETECTION OF
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Chairman: Associate Professor Mariana Nor Shamsudin, PhD

Faculty : Medicine and Health Sciences

Diarrhea is one of the leading causes of illnesses and death among children in developing countries, where an estimated 1.3 billion episodes and 4 to 10 million deaths occur each year in children below 5 years of age. The common pathogens of diarrhea are diarrheagenic *Escherichia coli* (DEC), Group A rotavirus, *Shigella* spp, *Salmonella* spp, *Campylobacter*, and *Vibrio cholerae*. Microbiological insights including phenotypic and genotypic characterisation are valuable approaches with application in management and prevention of diarrheal outbreaks by *E. coli*.

In the present study, the Random Amplified polymorphic DNA (RAPD) fingerprinting technique allowed genetic diversity assessment of 25 *E. coli* isolates. Six out of 20 arbitrary primers namely, OPAE 4, 9, 10, 11, 12 and 18 produced DNA fingerprinting patterns providing the discriminatory power and the display of the potential epidemiological and diagnostic markers. A highly significant finding from the DNA fingerprinting is the display of a predominant band at a size of 308 bp when arbitrary

fingerprinting is the display of a predominant band at a size of 308 bp when arbitrary OPAE-10 primer was used. The band consistently was amplified in *E.coli* strains from all sources but not in other Gram positive or Gram negative control strains.

The identity of this band after Blast analysis of TOPO 2.1–cloned sequence is a gene designated as *secD*. The finding is highly valuable with genotypic application as the gene is small in size and gave 100% homology to *E. coli* protein-export membrane, protein *secD*. In addition the high percentage of similarity (100%) of the RAPD-OPAE-10 marker to a fragment of *E. coli* genome emphasized the reliability of the marker as species-specific marker. The membrane assay developed using the probe designed from marker sequence consistently showed positive signal for all strains of *E. coli* tested but not for other Gram-positive and Gram-negative control strains tested, indicating the specificity and the sensitivity of *secD* marker primers and probes in species-specific detection of *E. coli* isolates. Although the RAPD OPAE-10 primers is commercially available, the RAPD OPAE-10 primer as *E. coli* diagnostic marker has not been reported previously. Another worthy finding from the RAPD study of the present work is the genetic diversity, and clonal groupings of *E.coli* strains from five different sources, determined simultaneously. The genetic diversity exhibited through the DNA fingerprinting patterns is accordingly observed in the antibiotic resistance pattern. A total of 70 isolates from different sources showed variations in resistance patterns to the 10 antibiotics tested from 61.2% *E. coli* isolates retrieved for antimicrobial resistance profiling. Overall, tetracycline and kanamycin were the most commonly reported antimicrobial agent resistance (81.0%) followed by chloramphenicol (76.0%), gentamycin (72.0%), and ampicillin (73.0%). While, resistances to ciprofloxacin of

24.0% norfloxacin, 27.0% and cefutixin, 40.0% were low in prevalence in all types of samples. The phenotypic based conventional biochemical identification method was performed in the study to confirm strain identity of all bacteria used in the research.

A genetic assessment of *E.coli* strain of various sources based on the *GAD* gene revealed another candidate gene for gene probe development. The 671bp gene has been reported to be homogeneously present in many *E. coli* strains but the present study illustrated the presences in *E.coli* strains of various sources, namely clinical, animal, sea water, river, and food. All *E. coli* isolates examined had the 671-bp and carried *GAD* gene. A membrane-based dot blot hybridization technique assay applying oligonucleotide probe designed from the *GAD* gene sequences was not only highly sensitive and specific for all *E. coli* isolates from different sources, but a rapid assay was optimized enabling the detection to be compacted in a very short time.

The genetic assessment study of *E.coli* strains enable the multiple PCR-based methods to be optimized for pathogenic strain determination. The contribution of six primer pairs specific for enterohemorrhagic *E. coli* (EHEC) *stx1* and *stx2* genes, enteropathogenic *E. coli* (EPEC) *bfp* and *eae* genes, enterotoxigenic *E. coli* (ETEC) *elt* gene, and enteroaggregative *E. coli* (EAEC) *EAST* gene can facilitate the rapid detection of these groups from different sources through a single tube PCR method.

In addition the virulence gene-based for pathogenic strains detection, another virulence determination Lipopolysaccharide (LPS) found in the Gram negative bacteria was applied for development of genera differentiation method. A simple biolysate-based

method for simultaneous detection and quantitation of the gram negative pathogenic bacteria based on lipopolysaccharide (LPS) component was optimized. The exposure of the marine animal lysate blood to the polysaccharides of gram negative pathogens results in activation of an intracellular coagulation. The hypothesis is the rate of coagulation is associated to the LPS chain length. Carbohydrate assay and turbidity assay are basis of the main methods used in this study. *E. coli*, *Samonella*. spp., and *V. cholerae* with similar initial cell count per ml had different absorbance readings by using the spectrophotometer and turbidity meter. The range of difference in absorbance is consistent between different selected gram negative genera. Two selected corresponding genes encoding the LPS are explored for future application as molecular detection assay. The *wzm* and *wzt* genes encoding O-polysaccharide genes were amplified in these pathogens and the LPS factor C were amplified from the marine lysate. The hybridization results clearly demonstrated that *wzm* gene, hybridized with anti-LPS factor C peptide of marine lysate, thereby; homogeneity of bacterial O-polysaccharide gene to factor C peptide could be exploited in genera differentiation method.

The significant impact of the study is the considerable advancement in the molecular protocols, molecular characterizations of local *E. coli* strains since protocols are highly optimized to suit local strains, and diverse *E. coli* sources were included in the study.

The present study established the framework for the potential improvement in methods using genotypic based approaches for sources tracing and related genes differentiation for identification and quantitation. The genomic based probe from the RAPD fingerprinting is notably worthy for *E.coli* species specific detection without PCR

approach and the newly developed epidemiological marker for *E. coli* from different sources is valuable in future molecular diagnostics of *E. coli*.

Abstrak tesis yang dikemukakan kepada Senat Universiti Putra Malaysia sebagai memenuhi keperluan untuk ijazah Doktor Falsafah

**PENAMBAHBAIKAN KAEDAH MOLECULAR UNTUK PENGESANAN
ESCHERICHIA COLI PATOGENIK**

Oleh

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Mei 2007

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Cirit-birit merupakan penyebab utama berlakunya penyakit dan kematian di kalangan kanak-kanak negara membangun, di mana sebanyak 1.3 billion episod dan 4 hingga 10 juta kematian dianggarkan berlaku setiap tahun dalam kanak-kanak di bawah umur 5 tahun. Patogen penyebab cirit-birit yang biasa adalah 'Diarrheagenic *Escherichia coli*' (DEC), rotavirus kumpulan A, spesis *Shigella*, spesis *Salmonella*, *Campylobacter*, dan *Vibrio cholerae*. Pandangan dari sudut mikrobiologi berdasarkan ciri-ciri fenotip dan genotip adalah cara pendekatan yang bernilai dalam mengurus dan menghindar daripada tersebarnya wabak cirit-birit yang berpunca daripada *E.coli*.

Dalam kajian ini, kaedah Randomly Amplified Polymorphic DNA (RAPD) digunakan untuk penilaian kepelbagaian genetik ke atas 25 isolat *E.coli* menggunakan 20 primer berbeza. Enam daripada 20 primer khususnya OPAE 4,9,10,11,12 dan 18 menghasilkan corak cap jari DNA dengan kuasa pembezaan yang menunjukkan potensi penanda ini

corak cap jari DNA dengan kuasa pembezaan yang menunjukkan potensi penanda ini sebagai penanda diagnostik dan epidemiologi. Penemuan yang sangat penting daripada corak jari DNA ini adalah pada saiz 308 bp yang mana jalur-jalur tebal dan terang hadir apabila primer OPAE-10 digunakan. Jalur-jalur tebal dan terang ini sentiasa hadir dalam semua 5 strain *E.coli* dari pelbagai sumber, namun tidak hadir dalam isolat Gram positif atau strain kawalan Gram negatif yang lain.

Identiti jalur ini selepas analisis BLAST dengan TOPO 2.1 ialah jujukan klon protein *secD*. Penemuan yang berharga daripada aplikasi genotip ini adalah saiz yang kecil di mana 100% homologi kepada membran pengangkut protein *E.coli* iaitu protein *secD*. Sebagai tambahan, peratus yang tinggi iaitu 100% persamaan dalam RAPD menggunakan primer OPAE-10 kepada genom *E.coli* menunjukkan bahawa penanda ini mampu digunakan sebagai penanda spesifik spesies. Di samping itu, esei membran yang dihasilkan menggunakan prob yang dicipta berdasarkan jujukan penanda menunjukkan signal positif untuk semua strain *E.coli* tetapi tidak pada isolat Gram positif dan Gram negatif yang lain, di mana ini membuktikan bahawa penanda *secD* adalah primer dan prob yang spesifik untuk mengesan spesies isolat *E.coli*. Walaupun primer RAPD OPAE-10 boleh didapati secara komersil, primer RAPD OPAE-10 sebagai penanda diagnostik *E.coli* masih belum dilaporkan dalam menjeniskan *E.coli* dari mana-mana sumber diagnostik. Sistem mengkelas RAPD yang digunakan dalam kajian ini merupakan yang pertama yang menyertakan 5 sumber berbeza serentak.

Penilaian genetik yang lain dalam mengesan strain *E.coli* dari pelbagai sumber adalah berdasarkan gen *GAD*. Ini menjadikan gen ini sebagai calon gen lain bagi kajian penghasilan prob gen. Gen pada saiz 671bp ini dilaporkan hadir secara homogenus pada

kebanyakan strain *E.coli* dari pelbagai sumber antaranya sumber klinikal, haiwan, air laut, sungai dan makanan. Kesemua isolat *E.coli* yang diperiksa membawa gen GAD pada 671bp. Esei membran menggunakan teknik hibridasi dot-blot mengaplikasikan prob oligonukleotida yang dicipta dari jujukan gen *GAD* bukan sahaja sangat sensitif dan spesifik ke atas semua isolat *E.coli*, malah ia merupakan esei yang cepat yang dihasilkan supaya dapat mengesan dalam masa yang singkat.

Penilaian kajian genetik ke atas strain *E.coli* membolehkan teknik berasaskan PCR dihasilkan dalam penentuan strain pembawa penyakit. Sumbangan enam pasang primer khas untuk gen-gen *stx1* dan *stx2* dari ‘enterohemorrhagic *E.coli*’ (EHEC), gen-gen *bfp* dan *eae* dari ‘enteropathogenic *E.coli*’ (EPEC), gen *elt* ‘enterotoxigenic *E.coli*’ (ETEC), dan gen *EAST* ‘enteroaggregative *E.coli*’ (EAEC) membantu dalam menghasilkan kaedah pengesanan pantas ke atas kumpulan *E.coli* dari pelbagai sumber dalam kaedah PCR tiub tunggal. Sebagai tambahan, kajian pengesanan berdasarkan gen virulen ke atas strain pembawa penyakit menggunakan Lipopolisakarida (LPS) yang didapati pada bakteria Gram negatif telah digunakan dalam menghasilkan kaedah genetik yang efektif.

Kaedah mudah berdasarkan hasil-pecahan biologi untuk pengesanan serentak bakteria patogen Gram negatif berdasarkan komponen lipopolisakarida telah diperbaiki. Pendedahan hasil-pecahan darah haiwan marin kepada polisakarida patogen Gram negatif mengaktifkan penggumpalan dalam sel. Hipotesis di sebalik kajian ini ialah kadar penggumpalan adalah berkaitan dengan panjang rangkaian LPS. Esei-esei karbohidrat dan kekeruhan merupakan kaedah utama dalam kajian ini. *E.coli*,

Salmonella spp., dan *Vibrio cholerae* dengan kiraan awal sel per mL yang sama mempunyai bacaan resapan yang berbeza tetapi konsisten dengan menggunakan spektrofotometer dan meter kekeruhan. Ciri molekul LPS yang berkait dipelopori untuk kegunaan masa depan dalam esei pengesanan gen peringkat molekul. Gen *wzm* dan *wzt* yang mengkodkan gen-gen O-polisakarida digandakan dalam patogen-patogen ini dan faktor C LPS digandakan daripada hasil-pecahan marin. Keputusan hibridasi dengan jelas menunjukkan gen *wzm* berhibridasi dengan peptida anti-LPS gen faktor C hasil-pecahan bio marin, di mana homogenesiti gen bakteria O-polisakarida kepada peptida factor C boleh dieksploitasi dalam kaedah pembezaan genera. Kesan ketara dari kajian ini adalah kemajuan yang agak pesat dalam protokol peringkat molekul, pencirian molekul strain-strain *E.coli* tempatan memandangkan protokol diperbaiki bersesuaian dengan strain-strain tempatan, dan *E.coli* pelbagai sumber dimasukkan dalam kajian ini.

Diversiti genetik yang dipamerkan dari corak cap jari DNA adalah berdasarkan corak kerentanan antibiotik, Tujuh puluh isolat *E.coli* dari pelbagai sumber menunjukkan corak kerentanan yang pelbagai kepada 10 ejen antimikrob yang diuji, 61.2% daripada isolat *E.coli* diambil semula untuk pemprofilan kerentanan antibiotik. Secara keseluruhan, tetracycline dan kanamycin merupakan agen antibiotik yang paling lazim dilaporkan iaitu sebanyak 81%, diikuti dengan chloramphenicol 76%, gentamycin 72%, dan ampicillin 73%. Sementara itu, kerentanan yang paling rendah dapat dilihat terhadap ciprofluoxacin (24.0%), norofluoxacin (27.0%), dan cefutixin (40.0%). Kaedah pengenalpastian biokimia konvensional berdasarkan fenotip dilakukan dalam kajian ini untuk mengesahkan identiti strain setiap bakteria yang digunakan dalam kajian ini.

Kesimpulannya, kajian ini telah mengukuhkan rangka kerja untuk memperbaiki potensi kaedah berasaskan genotip dalam mengesan sumber dan pembezaan gen berkaitan bagi tujuan pengenalanpastian dan kuantitasi. Prob berasaskan genom cap jari RAPD ini adalah penemuan penting yang bernilai ke atas pengesanan khusus isolat *E.coli* tanpa pendekatan PCR.

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I certify that an Examination Committee has on 24th May 2007 to conduct the final examination of Nagi Ahmed Abdullah AL-Haj on his Doctor of Philosophy thesis entitled “Improvement of Molecular Methods for Detection of Pathogenic *Escherichia coli*” in accordance with Universiti Pertanian Malaysia (Higher Degree) Act 1980 and Universiti Pertanian Malaysia (Higher Degree) Regulations 1981. The Committee recommends that the student be awarded the degree of Doctor of Philosophy.

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
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DECLARATION

I hereby declare that the thesis is based on my original work except for quotations and citations which have been duly acknowledged. I also declare that it has not been previously or concurrently submitted for any other degree at UPM or other institutions.



NAGI AHMED ABDULLAH AL-HAJ

Date: 26/08/08

TABLE OF CONTENTS

	Page
DEDICATION	ii
ABSTRACT	iii
ABSTRAK	viii
ACKNOWLEDGEMENTS	xiii
APPROVAL	xiv
DECLARATION	xvi
LIST OF TABLES	xxiv
LIST OF FIGURES	xxv
LIST OF ABBREVIATIONS	xxxi
 CHAPTER	
 I	
INTRODUCTION	1
Objectives	4
 II	
LITERATURE REVIEW	6
2.1 <i>Escherichia coli</i>	6
A) <i>E. coli</i> as water quality indicator	8
B) <i>E. coli</i> as Foodborne pathogens	8
2.2 Classification of Diarrheagenic <i>E. coli</i>	10
Enteropathogenic <i>E. coli</i> (EPEC).	10
Enterotoxigenic <i>E. coli</i> (ETEC)	11
Enteroinvasive <i>E. coli</i> (EIEC)	12
Enteropathogenic <i>E. coli</i> (EPEC)	13
Diffuse-adhering <i>E. coli</i> (DAEC)	14
Cytotoxic distending toxins- <i>E. coli</i> (CDT- EC)	14
Enterohemorrhagic <i>E. coli</i> (EHEC)	14
2.3 Laboratory diagnostics	16
Diarrheagenic <i>E. coli</i> growth requirements	16
Phenotypic study based on antibiotic resistance pattern	17
Genotypic methods	18
Polymerase Chain Reaction (PCR)	19

	Random Amplified Polymorphic DNA (RAPD)	21
	Multiplex Polymerase Chain Reaction (MPCR)	24
	Dot Blot Hybridization	25
	PCR Sequencing	27
2.4	Detection of Gram negative bacteria based on conventional and molecular methods	28
III		31
	MATERIALS AND METHODS	
3.1	Bacterial sources	31
3.1.1	Phenotypic characteristics tests for <i>E. coli</i> identification	35
	A) Growth on MacConkey agar	35
	B) Growth on Eosin Methylene Blue Agar (EMB)	35
	C) Growth on Chromocult Coliform Agar (CCA)	35
	D) Gram Staining	37
	E) Oxidase Test	36
	F) Catalase Test	36
	G) API 20E	37
	H) Isolation of <i>E. coli</i> from different water sources	37
	I) Food samples processing.	38
3.1.2	Phenotypic method for characterization of n antibiotic resistance	39
	A) Determine of diversity of <i>E. coli</i> based on antibiotic resistance patterns among isolates of different sources	39
	B) Data Analysis.	40
3.2	Genotypic based methods for characterization of bacterial isolates: RAPD fingerprinting	40
3.2.1	Clonal diversity determination <i>E. coli</i> species-specific marker screening from RAPD fingerprint	40
	A) Propagation of Bacterial Culture	40
	B) Total DNA Extraction	41
	C) Quantitation of DNA	42
	D) Agarose Gel Electrophoresis of genomic DNA	42
	E) PCR for DNA finger printing	43

I) RAPD Analysis	44
II) Identification of diagnostic marker from the RAPD-PCR Fingerprint	45
III) Gel purification of the RAPD Specific Band	45
IV) Cloning of RAPD Specific Band	46
V) One shot transformation reaction	46
VI) Analysis of positive clones	47
VII) Extraction of plasmid	47
VIII) Restriction Enzyme Analysis of 2.1-TOPO Clones	48
IX) DNA Sequencing	48
X) Dot blot assay for detection of a rapid-based marker (<i>secD</i> gene)	49
a. Preparation of labeled probe	49
b. Blotting the samples on the membrane	49
c. Immobilization	50
d. Hybridization	50
e. Chemiluminescent Detection	50
3.3 Characterization using GAD gene	52
3.3.1 Amplification of <i>GAD</i> gene	52
A. Development of a probe based on <i>Gad</i> gene sequence for the identification of <i>E. coli</i> from different sources	53
B. DNA sequencing and analysis of <i>GAD</i> gene sequence from different sources of <i>E. coli</i>	54
3.4. Differentiation of diarrheagenic <i>E. coli</i> from various sources based on virulent gene	55
A) Designing of primers specific for different virulent genes	55
B) PCR Optimization for detection of virulent genes	57
I. Optimization of cocktail mixture.	57
II. Optimization of PCR cycling conditions	58
III . Optimization of Qigen kit extraction DNA	58
IV . PCR assay employing all optimized	59
3.5 Development of Lipopolysaccharide (LPS)-based assay for	59

detection of <i>E. coli</i> and related genera	
A) Biolysate-based detection of <i>E. coli</i>	60
B) Biolysate-based turbidity assay	60
C) Carbohydrate assay	60
3.5.1 Carbohydrate and turbidity assays for detection and quantitation	61
A) Direct plating and enrichment	61
Genotypic detection of gram negative bacteria through dot blot	62
hybridization of LPS from <i>E. coli</i> and target peptide gene from	
horse shoe crab	
A) DNA extraction of <i>Factor C</i> marine biolysate	62
B) Detection and Amplification of Factor C anti-LPS gene of	63
marine biolysate	
C) DNA Extraction of LPS biosynthesis gene from <i>E. coli</i> , <i>V.</i>	64
<i>cholera</i> and <i>Salmonella</i> spp	
D) Detection and amplification of <i>wzm</i> , <i>wzt</i> from <i>E. coli</i> , <i>V. cholera</i>	65
and <i>Salmonella</i> spp	
E) PCR Product Purification	66
F) Gene sequencing of <i>Factor C</i> of marine biolysate gene and LPS	67
genes of <i>E. coli</i> , <i>V. cholera</i> and <i>Salmonella</i> spp	
G) Dot blot hybridization	68
IV	69
RESULTS	
4.1 Characterization Tests	69
A) Growth on MacConkey agar	69
B) Growth on Eosin Methylene blue agar (EMB)	69
C) Growth on Chromocult Coliform Agar (CCA)	69
D) Gram stain	71
E) Catalase test	70
F) Oxidase test	70
G) API 20E	70
4.1.2 Phenotypic based methods for characterization antibiotic	73

	susceptibility characterization	
	A) Determination diversity of <i>E. coli</i> based on antibiotic resistance patterns among isolates of different sources	73
	Genomic Extraction	81
4.2	Genotypic based methods for characterization of bacterial samples: RAPD fingerprinting	84
4.2.1	Screen for <i>E. coli</i> species-specific marker from RAPD fingerprint	84
	RAPD analysis	93
	Genetic distance and percentage of similarity	95
	Dendrogram	93
	II) Identification of diagnostic marker from the RAPD-PCR fingerprint	97
	Cloning in plasmid vectors	97
	Analysis of clones	99
	DNA sequencing of diagnostic marker from the RAPD-PCR Fingerprint	99
	Amplification of <i>secD</i> gene from <i>E. coli</i> and <i>Salmonella</i> spp	104
	Dot blot assay for detection of a rapid-based marker <i>secD</i> gene	104
4.3	Characterization of <i>E. coli</i> isolates from different sources using <i>GAD</i> gene	106
4.3.1	Amplifying <i>GAD</i> gene from different groups of <i>E. coli</i>	106
4.3.2	DNA sequencing of <i>E. coli</i> specific gene from different sources	108
	Development of a probe based on <i>gad</i> gene sequence for the identification of <i>E. coli</i> from different sources	116
4.4.	Differentiation of diarrheagenic <i>E. coli</i> of various sources based on virulent gene	118
	a) Design of primers specific for different virulent genes	18
	b) Amplification of <i>bfp</i> gene (Bundle forming pilus) from different sources	123
	c) Sequence analysis of genomic <i>bfp</i> gene	123
	d) Detecting of virulent genes of diarrheagenic <i>E. coli</i> by multiplex	127

	PCR	
4.5	Developing lipopolysaccharide-based (LPS) assay for detection of <i>E. coli</i> and related genera	129
	A) Biolysate-based turbidity assay	129
	B) Carbohydrate based assay	130
	Molecular detection of Gram negative bacteria through dot blot hybridization of LPS from <i>E. coli</i> and target peptide gene from horse shoe crab	132
	a) DNA extraction of <i>Factor C</i> marine biolysate	132
	b. Detection and amplification of Factor C anti-LPS gene of marine <i>limulus ameobocyte</i>	132
	i) Sequence analysis of <i>Factor C anti-LPS</i> gene (<i>Tachypleus tridentatus</i> anti-lipopolysaccharide factor) anti-LPS	134
	c. DNA Extraction of LPS biosynthesis gene from <i>E. coli</i> , <i>V. cholera</i> and <i>Salmonella</i> spp	134
	d. Detection and amplification of <i>wzm</i> and <i>wzt</i> from <i>E. coli</i> , <i>V. cholera</i> and <i>Salmonella</i> spp	134
	e. Detection and Amplification of <i>wzm</i> and <i>wzt</i> from <i>E. coli</i> , <i>V. cholera</i> and <i>Salmonella</i> spp	135
	f. Sequence analysis of <i>wzm</i> gene from <i>E. coli</i> , <i>Salmonella</i> spp and <i>V. cholerae</i> (<i>Elongation factor Tu</i>)	135
	h. Sequence analysis of <i>wzt</i> gene from <i>E. coli</i> , <i>Salmonella</i> spp and <i>V. cholera</i>	136
	Dot blot hybridization	146
V		151
	DISCUSSION	
5.1	Phenotypic approaches using conventional culture methods	151
5.2.	Molecular based methods for characterization of bacterial isolates: RAPD fingerprinting for diversity and markers determination.	160
5.2.1	DNA sequencing of diagnostic marker from the RAPD-PCR	170

	fingerprint	
5.2.2	Dot blot assay for detection of a rapid-based marker (<i>secD</i> gene)	172
5.3.	Characterization of <i>E. coli</i> isolates from different sources using <i>GAD</i> gene	173
5.3.1	Amplification <i>GAD</i> gene in strains from different groups of <i>E. coli</i>	173
5.3.2	DNA sequencing of <i>E. coli</i> specific gene from different sources	175
	Development of a probe based on <i>gad</i> gene sequence for the identification of <i>E. coli</i> from different sources through dot blot hybridization assay	176
5.4	Differentiation diarrheagenic <i>E. coli</i> of various sources based on virulent gene	179
	a) Designing of primers specific for different virulent genes	179
	b) Characterization of <i>bfp</i> gene from different sources	181
5.4.1	Molecular method for detecting virulence genes of diarrheagenic <i>E. coli</i> by multiplex PCR	183
5.5	Development of Lipopolysaccharide (LPS)-based assay for detection of <i>E. coli</i> and related genera	186
	Detection of Gram negative bacteria through dot blot hybridization of LPS from <i>E. coli</i> and a target peptide gene from horse shoe crab	192
	a) Detection and amplification of <i>wzm</i> , <i>wzt</i> from <i>E. coli</i> , <i>V. cholera</i> and <i>Salmonella</i> spp	193
	b) Dot blot hybridization	195
VI	CONCLUSION	199
VII	REFERENCES	204
	APPENDICES	237
	BIODATA OF THE AUTHOR	257
	PUBLICATIONS	258

LIST OF TABLES

Table	Page
1 Sources of <i>E. coli</i> isolates used in this study	31
2 Typed of <i>E. coli</i> used as reference strain	34
3 Control strain of Gram negative and gram positive genera	34
4 Sequences of primers screened for RAPD procedure	43
5 Designed primer used for amplification of the <i>secD</i> gene	52
6 Designed primer and probe used for amplification and detection of the GAD gene	53
7 Designed primers used in the multiplex PCR for detection of diarrheagenic groups of <i>E. coli</i>	56
8 Primer sequence of Factor C (<i>anti-LPS</i>) gene	63
9 The sequences of primers and biotin labeled probes used in detection and amplification of <i>wzm</i> and <i>wzt</i> genes used in this study	67
10 Rate antimicrobial resistance among of <i>E. coli</i> isolates	73
11 Percentage of antimicrobial resistance in <i>E. coli</i> isolated from various sources (Human, sea water, river water, foods, and animals)	74
12 Rates of multidrugs resistance among <i>E. coli</i> isolates of various sources	81
13 Value of genetic distance (below diagonal) and percentage of similarity (above diagonal) among <i>E. coli</i> isolates from varieties sources 1-5 Human, 6-10 sea water, 11-15, River water, 16-20 food, 21-25 animal based on RAPD data and, using the computed formula of Nei and Li.	95
14 Virulence genes distribution of <i>E. coli</i> isolates from five different sources amplified by PCR	119